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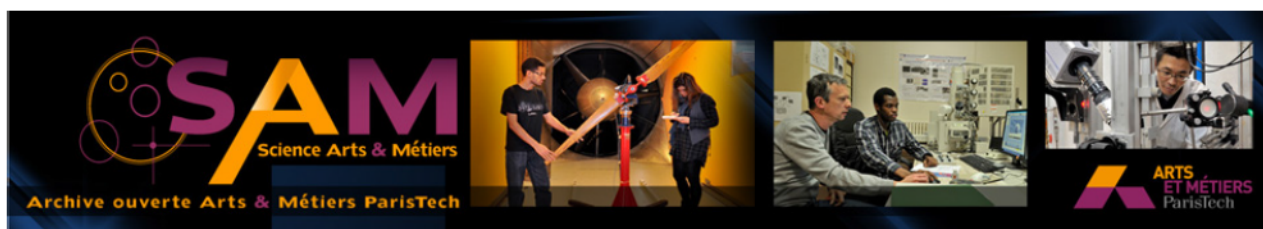
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# Validation of 3D spino-pelvic muscle reconstructions based on dedicated MRI sequences for fat-water quantification

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## Abstract

**Objectives.** – To evaluate a protocol, including MRI acquisition with dedicated sequences for fat-water quantification and semi-automatic segmentation, for 3D geometry measurement and fat infiltration of key muscles of the spino-pelvic complex.

**Materials and methods.** – MRI protocol: two axial acquisitions from the thoraco-lumbar region to the patella were obtained: one T1-weighted and one based on the Dixon method, permitted to evaluate the proportion of fat inside each muscle. Muscle reconstruction: with Muscl'X software, 3D reconstructions of 18 muscles or groups of muscles were obtained identifying their contours on a limited number of axial images 3D references were obtained only on T1 acquisitions identifying the contour of the muscles on all axial images. Evaluation: for two volunteers, three operators completed reconstructions three times across three sessions. Each reconstruction was projected on the reference to calculate the 'point to surface' error. Mean and maximal axial section, muscle volume, and muscle length calculated from the reconstructions were compared to reference values, and intra- and inter-operator variability for those parameters were evaluated.

**Results.** – 2xRMS 'point to surface' error was below 3 mm, on average. The agreement between the two methods was variable between muscles [–4.50; 8.00%] for the mean axial section, the length and the volume. Intra- and inter-operator variability were less than 5% and comparison of variability for the Fat and T1 reconstructions did not reveal any significant differences.

**Discussion.** – Excellent inter- and intra-operator reliability was demonstrated for 3D muscular reconstruction using the DPSO method and Dixon images that allowed generation of patient-specific musculoskeletal models.

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## 1. Introduction

The muscular system plays an essential role in the maintenance of postural balance; however, in clinical practice as well as research, investigation into the relationship between the muscular system and postural pathologies such as adult spinal deformity, has been limited.

The current lack of knowledge related to soft tissue stabilizers may be attributed to the absence of a relevant tool to evaluate the muscular system as a whole. The large number of muscles involved in postural maintenance makes a global analysis difficult and time consuming. In the case of adult spinal deformity (ASD), recent research has highlighted the critical role of sagittal spino-pelvic alignment in patient-reported pain and disability

[1–5]. Therefore, key muscles involved in pelvic-positioning and lumbar-stabilization are at the forefront of research needs.

Preliminary efforts have been directed towards understanding the muscular envelope of the spine using histological analyses [6–9], measurement of muscular strength [10–12], and measurement of electromyographical signals [13,14]. However, these approaches are not adapted for study of a large number of muscle groups. Other studies, using imaging such as MRI or CT scan, correlated measurement of the muscle cross-sectional areas (via ultrasound, CT-scan or MRI) [15–23], or measurement of muscular density (via CT-scan or MRI) [10–12,17–21,23–25], with chronic back pain or spinal surgery outcome. The limitations of these approaches lie in the difficulty to represent variability in volume of an entire muscle [26].

To overcome these limitations, Jolivet et al. [27,28] developed a method of three-dimensional muscle reconstruction via segmentation of a small number of axial images (MRI or CT-scan). This method, based on the deformation of a parametric

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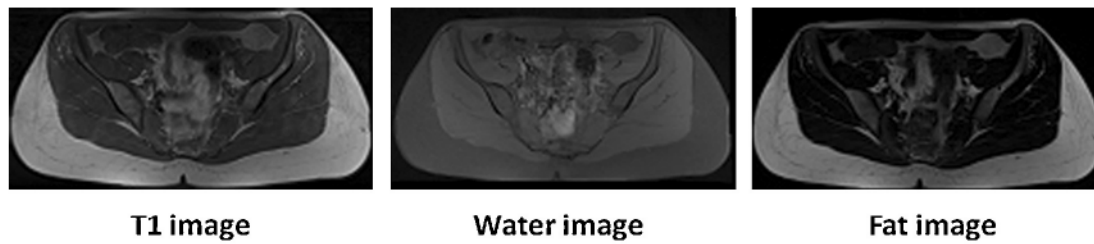


Fig. 1. Examples of a T1 (left), water (center), and Fat (right) image.

specific object (DPSO), has been successfully implemented with CT scans for analysis of muscles involved in knee motion [29] and muscle groups around the hip joint [28]. The CT-scan modality presents two advantages: good contrast quality for muscle segmentation and good reliability in terms of fat and muscle density. CT scans allow for both a reproducible analysis of muscle geometry and a quantified evaluation of fat infiltration [28]. Nevertheless, the radiation exposure from CT scans renders it unacceptable as a tool for studies involving ASD patients who are already frequently subjected to radiographic examination. Notably, the DPSO method has also been performed using MRI T1 sequences [29]. However, the inhomogeneity in the magnetic field applied did not allow an accurate quantification of the fat infiltration (without fat infiltration, the muscle volume is an incomplete descriptor).

In order to avoid the problem of inhomogeneity and obtain an accurate quantification of fat infiltration, Dixon et al. [30] developed a specific acquisition sequence where two images are obtained: one in which the intensity of each voxel is correlated with the quantity of fat and the other in which the intensity of each voxel is correlated with the quantity of water. This method was then improved by Glover et al. [31]. To our knowledge, the feasibility of 3D muscular reconstruction on MRI with the Dixon method was not studied.

The objective of this study was to evaluate the feasibility of an MRI protocol with dedicated sequences for fat-water quantification to assess the 3D geometry and homogeneity (fat infiltration) of key muscles in the spino-pelvic complex.

## 2. Methods

### 2.1. Subject sample

Two asymptomatic female adult volunteers were included in this pilot study: volunteer A (35 years, 68 kg) and volunteer B (38 years, 91 kg).

### 2.2. MRI acquisition

MRI was performed on a 3T whole-body scanner (Magnetom Verio, Siemens Healthcare, Erlangen, Germany) using a 24-channel spine matrix coil and three 4-channel flex coils from the same vendor. The imaging protocol included a T1-weighted turbo spin-echo (T1 TSE) sequence (TR/TE = 1220/11 ms, acquisition matrix =  $512 \times 384$ , in plane resolution =  $0.98 \times 0.98 \text{ mm}^2$ , slice thickness = 5 mm, slice

gap = 5 mm, parallel imaging acceleration factor (iPat) = 2, 40 slices, flip angle =  $150^\circ$ , bandwidth = 219 Hz/pixel, turbo factor = 5, acquisition time = 2:15 min/40 slices, 160 slices by patients) and a T1-weighted TSE sequence for applying the three point Dixon method [30–33] (TR/TE = 829/15.7 ms, acquisition matrix =  $512 \times 384$ , in plane resolution =  $0.98 \times 0.98 \text{ mm}^2$ , slice thickness = 5 mm, slice gap = 5 mm, iPat = 2, 40 slices, flip angle =  $150^\circ$ , bandwidth = 315 Hz/pixel, turbo factor = 3, echo spacing = 15.7, acquisition time = 4:38 min/40 slices, 160 slices by patients). Water and Fat images were automatically generated by the scanner from the TSE images for the three point Dixon method (Fig. 1). Both sequences had exactly the same slice position and orientation. Image volume covered the proximal tibia to the lumbar spine (Th12 vertebra) and was acquired in three stages. Total acquisition time was 45 min.

### 2.3. 3D muscle reconstruction (DPSO method)

The 3D reconstruction of individual muscles was performed using Muscl’X software, a custom software, (Laboratory of Biomechanics, Arts et Métiers ParisTech, France). Using the axial MR images, the software generated the 3D geometry of each muscle. The reconstruction technique was based on the DPSO algorithm as described in the literature [27–29] and briefly summarized hereafter (Fig. 2).

For each muscle, a subset of MRI axial slices (MSS: manually segmented slices) were manually segmented. The optimal percentage of MSS (defined as the number of MSS divided by the total number of slices covering the entire muscle) of each muscle is reported in Table 1 [34]. Of note, complex muscle geometries required a larger percentage of MSS than simple geometries. Using contrast differences, these manual segmentations were then optimized (Fig. 2a). The contours were then approximated by ellipses (Fig. 2b) and cubic spline interpolation was used to interpolate ellipses in all non-outlined slices covering the muscle (Fig. 2c). These interpolated ellipses generated a 3D parametric object (Fig. 2d). Finally, using a kriging algorithm [35], the parametric object was deformed non-linearly using the manual segmentations of MSS as control points (Fig. 2e). Contrast enhancements were used to optimize the segmentation at each slice. Once all muscles were reconstructed, a geometry-correction algorithm was applied to eliminate muscle segmentation interpenetration and a visual verification was performed to correct local geometric deformities. Finally, 3D meshed reconstructions of each muscle were obtained.



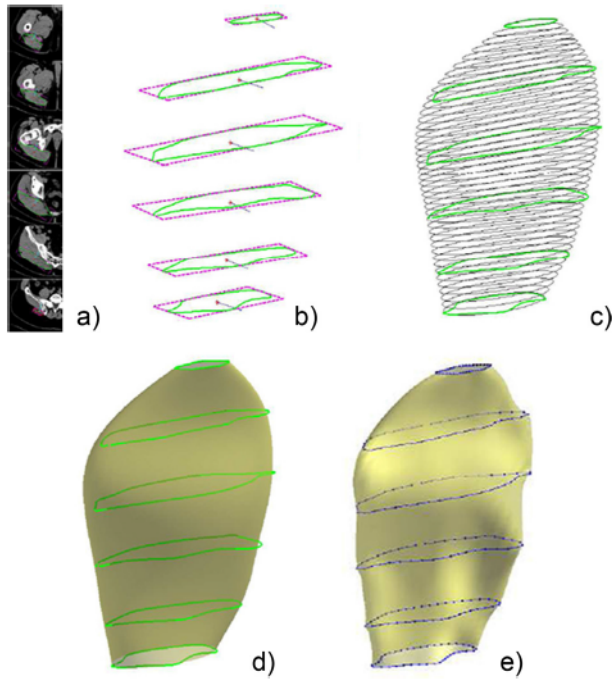


Fig. 2. Overview of the reconstruction method [28]: a: subset of MRI axial slices with manually segmented axial sections; b: contours approximated by ellipses [represented on the figure by a green rectangle whose length and width correspond to the major and minor axes of the ellipses]; c: cubic spline interpolation used to interpolate ellipses in all non-outlined slices covering the muscle; d: interpolated ellipses generated a 3D parametric object; e: non-linear deformation of the parametric object was deformed using the manual segmentations.

#### 2.4. Muscles of interest

Table 1 describes the right- and left-sided muscles or groups of muscles analyzed in this study. Muscles were chosen based on their potential role in regulating the position of the pelvis and the spine. Because the delineation of certain muscles was difficult and in order to decrease the number of reconstructed muscles,

Table 1  
Muscles analyzed in this study, and optimal percentage of MSS slices.

Muscle reconstructed(right and left)	Percentage of MSS
Adductor	20
Biceps femoris	12
Erector spinae	15
Gluteus maximus	18
Gluteus medius	25
Gluteus minimus	30
Gracilis	10
Iliacus	25
Obliquus	20
Psoas	10
Quadratus lumborum	18
Rectus abdominus	12
Rectus femoris	13
Sartorius	10
Semi-membranous tendinosis	11
Tensor Fascia Lata	15
Vastus lateralis inter	15
Vastus medialis	15

MSS: manually segmented slices.

some of the muscles were regrouped. The adductor longus, brevis and magnus were reconstructed into a single group named “Adductor”. In addition, the transversus abdominis muscle, internus obliquus, and externus obliquus were considered as a single group named “Obliquus” (they were reconstructed from their caudal insertion up to the liver). The rectus abdominus, the psoas, and the erector spinae were reconstructed from their caudal insertion to the superior endplate of the first lumbar vertebra. The short- and long- heads of the biceps femoris were grouped together (“Biceps femoris”), as were the semimembranosus and semitendinosus muscles (“Semi-membranous tendinosus”), and the vastus lateralis and vastus intermedius muscles (“vastus lateralis inter”). In total, 18 muscles, right and left, were analyzed.

#### 2.5. MR parameters

Based on 3D reconstructions, the following parameters were calculated for each muscle: mean and maximal axial section (AS), muscle volume, and muscle length. Muscle length was defined as the sum of distances between barycenters of all consecutive slices.

From the Water and Fat images, the relative fat content for each voxel, fat/water ratio, was calculated using Eq. (1).

$$\text{Fat/Water Ratio} = 100 \times \frac{\text{SI}_{\text{fat}}}{\text{SI}_{\text{fat}} + \text{SI}_{\text{water}}} \quad (1)$$

where  $\text{SI}_{\text{fat}}$  represents the signal intensity of the Fat image and  $\text{SI}_{\text{water}}$  represents the signal intensity of the Water image. For each muscle, all voxels contained in the muscle outline were identified. The average fat/water ratio was calculated over all voxels contained in each muscle allowing a quantification of the fat inside each muscle.

#### 2.6. Evaluation of the protocol

All the operators involved in the protocol evaluation were experienced in reading muscular anatomy on MR images and received training in the use of the software (Muscl’X). For the DPSO method, the first and last slices of each muscle were manually segmented and used as limits for all reconstructions; all operators used exactly the same number of MSS (Table 1) while the selection of the actual MSS was left to the operator’s discretion.

#### 2.7. Evaluation of the DPSO method

For each muscle of the studied volunteers, a reference object (i.e. 3D geometry) was generated on T1 images by manually contouring all MR axial images covering it. Fig. 3 presents the 3D reconstruction of volunteer A. This method of reconstruction was named the reference method and the 3D reconstruction objects were named references. The references for volunteers A and B were obtained by one operator. Using the DPSO method and the number of MSS presented in Table 1, this operator reconstructed once on the Fat images and once on the T1 images all the muscles for both volunteers.

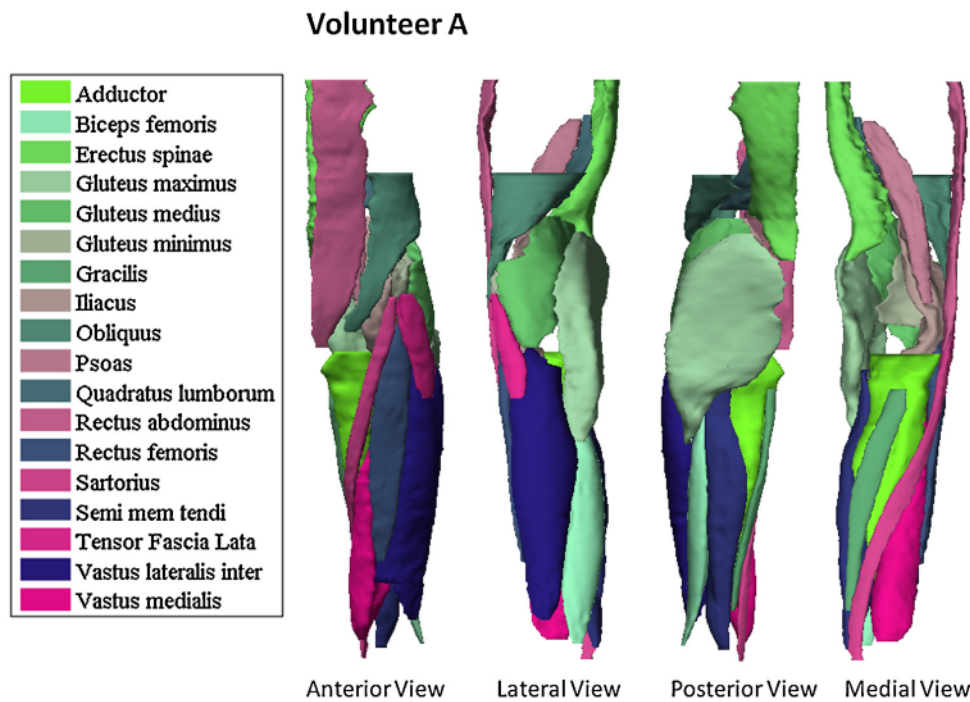


Fig. 3. Reference reconstruction for volunteer A including only left muscles; anterior, lateral, posterior and medial views.

The agreement between the DPSO with T1 or Fat images and the reference method (T1 images) was assessed with the percentage of difference between the methods [36] for the following parameters: maximal AS, mean AS, length, and volume. The mean, standard deviation (STD), maximum and minimum amount of difference between the reference values and the reconstruction values were expressed as a percentage of the reference values. The limits of the agreement were defined by the interval  $[\text{Mean} - 2 \times \text{STD}, \text{Mean} + 2 \times \text{STD}]$ .

## 2.8. Reproducibility of the DPSO method for T1 and Fat images

For this study, the intra- and inter-operator variability of the maximal AS, mean AS, length and volume were calculated in addition to shape accuracy. In this context, three operators using the DPSO methods completed the reconstruction of each muscle for each volunteer three times based on the T1 images ("T1 reconstruction") and three times based on the Fat images ("Fat reconstruction"). Thus, for each muscle of each volunteer, reconstruction was performed nine times for each sequence (3 operators  $\times$  3 sessions for T1 and Fat images). The muscular reconstructions for each volunteer were repeated with a minimum interval of 3 days for each operator.

### 2.8.1. Intra- and inter-variability

For the maximal AS, mean AS, length and volume, the reliability (intra-operator variability) and the reproducibility (inter-operator variability) from the T1 and Fat reconstructions were computed according to the ISO standard 5725-2:1994 [37] as the coefficient of variation expressed as a percentage. Two

different coefficients of variation were calculated: the coefficient of variation of each operator for each volunteer and the coefficient of variation of the average of the three operators for each volunteer. The intra-operator variability was the root mean square of the first coefficient of variation and the inter-operator variability was the root mean square of the sum of the two coefficients of variation. The intra- and inter-operator variability for the average fat/water ratio was calculated only using the Fat reconstructions.

### 2.8.2. Shape accuracy

In order to evaluate shape accuracy of the reconstructions, the data obtained from the three operators were compared to the reference previously reconstructed for the evaluation of the DPSO method. Differences in shape between the reference and each reconstruction obtained with the DPSO method (T1 and Fat reconstructions) were evaluated by projecting perpendicularly to the reference in three dimensions the points of the reconstructed muscles onto the reference surface. The 95% confidence interval for the average point-to-surface-distance [38] was evaluated as two times the root mean square (2xRMS). The mean, standard deviation, minimum and maximum values of the 2xRMS distances were calculated separately for the T1 reconstructions and the Fat reconstructions pooling the reconstructions of all operators together.

Statistical analyses were conducted using the software SPSS version 17 (SPSS, Inc, Chicago, IL). Differences between the reproducibility of T1 and Fat reconstructions were investigated using a two-sided paired *t*-test with a level of significance of 0.05.



Table 2

Results for volunteers A and B obtained with reference method (except fat/water ratio: average on all the Fat reconstructions).

Name	Max AS (mm <sup>2</sup> )		Mean AS (mm <sup>2</sup> )		Length (mm)		Volume (mm <sup>3</sup> )		Fat/water ratio (%)	
	A	B	A	B	A	B	A	B	A	B
Adductor L	4810	5426	2845	3050	300	329	828738	932146	12.78	20.45
Adductor R	4865	5173	2714	2990	301	327	776222	916293	12.45	18.11
Biceps femoris L	1366	1438	662	705	328	357	213860	238231	14.03	23.33
Biceps femoris R	1265	1428	632	717	337	352	207257	245682	17.76	21.74
Erectus spinae L	2533	2768	1889	1832	267	212	438288	341158	21.90	28.30
Erectus spinae R	2714	2615	1890	1830	261	212	439112	340376	23.27	28.49
Gluteus maximus L	4273	5943	2690	3321	300	318	737806	892654	18.75	33.84
Gluteus maximus R	3963	5561	2571	3154	289	319	692013	878789	19.63	33.70
Gluteus medius L	3054	3040	1744	1846	205	205	303772	302643	12.41	19.27
Gluteus medius R	2728	3040	1653	1673	210	206	287706	298895	12.72	19.12
Gluteus minimus L	1362	1195	943	804	131	116	93116	70784	15.60	19.54
Gluteus minimus R	1647	1161	976	689	133	125	101465	63928	16.71	18.25
Gracilis L	331	544	229	351	300	301	65342	98798	12.05	20.59
Gracilis R	367	592	245	396	297	288	69990	107238	10.99	19.36
Iliacus L	1317	1263	877	766	253	246	200654	163212	15.59	21.67
Iliacus R	1336	1341	858	745	245	242	187503	158666	12.82	17.53
Obliquus L	3046	2537	1157	1228	245	213	234643	225859	25.85	31.44
Obliquus R	3032	2315	1156	1150	263	210	246104	212022	25.00	33.22
Psoas L	1325	1155	721	657	258	237	171766	146658	17.20	26.03
Psoas R	1356	1185	740	654	255	240	176296	146012	17.03	23.91
Quadratus lumborum L	862	536	459	365	153	137	55572	42194	18.02	29.33
Quadratus lumborum R	701	567	347	363	159	139	45449	41563	22.20	26.60
Rectus abdominus L	1069	729	657	612	357	303	226174	178018	24.54	32.28
Rectus abdominus R	816	798	623	594	359	301	214962	172540	22.88	31.46
Rectus femoris L	878	1196	526	700	309	327	161707	222483	13.20	15.60
Rectus femoris R	934	1075	546	641	304	326	162332	203609	10.37	15.49
Sartorius L	330	438	214	308	498	465	99095	132455	18.25	23.86
Sartorius R	308	406	234	317	470	468	100298	134812	14.02	22.94
Semi mem tendi L	1575	2416	906	1343	370	367	328975	474250	18.78	21.68
Semi mem tendi R	1425	2358	906	1283	372	378	329319	459471	16.14	21.48
Tensor Fascia Lata L	596	672	378	425	144	151	52234	60726	16.78	22.75
Tensor Fascia Lata R	567	632	369	427	141	148	50795	58669	17.02	23.96
Vastus lateralis inter L	3648	4558	2288	2657	371	372	810584	940247	11.11	16.37
Vastus lateralis inter R	3608	4675	2212	2762	383	372	794354	964067	11.40	15.16
Vastus medialis L	1837	2235	1003	1194	354	322	344227	373751	10.84	15.10
Vastus medialis R	1598	2123	956	1290	331	308	299664	384815	9.49	13.24

AS: axial section.

### 3. Results

#### 3.1. Reconstruction time

The time to obtain a reconstruction (including visual verification to correct local geometric errors) of the muscles for one volunteer was 7 hours. For the reference, the time of reconstruction was between 14 and 15 hours.

#### 3.2. Individual results per subject

The reference values for volunteers A and B are presented in Table 2. This table includes results of the maximal AS, mean AS, length, volume and fat/water ratio (average of all Fat reconstructions) for each muscle.

#### 3.3. Evaluation of the DPSO method

Table 3 illustrates the agreement between the DPSO methods with Fat and T1 images and the reference methods. The

limits of agreement between the DPSO methods with Fat and the reference methods were  $[-3.50; 7.89]$  for the volume. For both sequences, the maximal AS showed the largest errors. The results demonstrated that for the volume and the mean AS, the agreement between the DPSO method with T1 images and the reference method was better than the agreement between the DPSO method with Fat images and the Reference. On average, the DPSO method, as compared to the reference method, tended to underestimate all parameters (between 0.74% to 2.20%).

#### 3.4. Intra- and inter-operator variability (DPSO method with T1 and Fat images only)

The intra- and inter-operator variability results for the T1 and Fat reconstructions of each muscle are summarized in terms of mean, standard deviation, maximal and minimal values for all the muscles in Tables 4 and 5. On average, the intra- and inter-operator variability was less than 5% for all parameters. For both T1 and Fat images, only the gluteus minimus (both left and right)

Table 3  
Variability in percentage between reference methods and DSPO methods for T1 and Fat images.

On all the muscles and both volunteers	100 × (reference method – DSPO Method)/reference method for one operator and both volunteers							
	Max AS		Mean AS <sup>a</sup>		Length		Volume <sup>a</sup>	
	T1	Fat	T1	Fat	T1	Fat	T1	Fat
Mean	1.21	1.16	0.74	2.20	0.91	0.44	1.10	2.19
Std	4.87	6.17	2.57	2.82	1.79	2.44	2.50	2.85
Limits of agreement sup (Mean + 2 × Std)	10.95	13.49	5.87	7.84	4.49	5.32	6.10	7.89
Limits of agreement inf (Mean – 2 × Std)	–8.53	–11.18	–4.39	–3.45	–2.68	–4.45	–3.91	–3.50
Max	15.43	15.57	6.01	6.92	6.18	10.39	5.92	6.92
Min	–9.41	–17.75	–6.22	–6.50	–2.96	–4.08	–6.62	–6.46

AS: axial section.

<sup>a</sup> Significant difference between Fat and T1.

Table 4  
Intra-operator variability in percentage (coefficient of variation).

	Max AS		Mean AS		Length <sup>a</sup>		Volume		Fat/water ratio
	T1	Fat	T1	Fat	T1	Fat	T1	Fat	Fat
Mean(on all the muscles)	4.08	3.76	2.16	2.05	1.78	1.26	2.16	2.05	2.23
Std (on all the muscles)	2.54	2.19	0.87	1.02	2.05	1.02	0.86	1.01	1.25
Max (on all the muscles)	12.49	10.66	3.81	5.60	9.93	5.16	3.75	5.62	5.56
Min (on all the muscles)	0.93	1.44	0.94	0.89	0.17	0.10	0.95	0.90	0.82

AS: axial section.

<sup>a</sup> Significant difference between Fat and T1.

Table 5  
Inter-operator variability in percentage (coefficient of variation).

	Max AS		Mean AS		Length <sup>a</sup>		Volume		Fat/water ratio
	T1	Fat	T1	Fat	T1	Fat	T1	Fat	Fat
Mean(on all the muscles)	4.63	4.50	2.56	2.61	1.86	1.42	2.55	2.61	2.98
Std (on all the muscles)	3.12	3.33	1.09	1.82	2.10	1.27	1.07	1.82	1.76
Max (on all the muscles)	15.65	19.62	5.80	11.67	9.93	5.93	5.78	11.80	10.59
Min (on all the muscles)	1.01	1.46	0.94	0.93	0.17	0.14	0.95	0.93	1.24

AS: axial section.

<sup>a</sup> Significant difference between Fat and T1.

had an intra-operator variability greater than 5% for the mean AS, length, and volume.

The other muscles with an inter-operator variability greater than 5% were the rectus abdominus and the gluteus medius. For the T1 image, the inter-operator variability of the length was equal to 5.87% for the gluteus medius right. On T1 images, the rectus abdominus left inter-operator variability was 5.80% for the mean AS and 5.78% for the volume.

For the average of fat/water ratio, the left and right gluteus minima and the left vastus medialis were the only muscles with an intra-operator variability greater than 5%. Except for the length, no significant differences were found between T1 and Fat reconstructions in terms of intra- or inter-operator variability.

### 3.5. Shape accuracy analysis

The results of the point-to-surface-distances for the T1 and Fat reconstructions of each muscle are summarized in terms

of mean, standard deviation, and minimal and maximal values for all muscles in Table 6. The mean 2xRMS values were less than 3 mm (i.e. less than 3 voxels), with significantly smaller values for the T1 reconstructions than for the Fat reconstructions. The minimum 2xRMS was also significantly smaller for the T1 images. The maximal 2xRMS value in terms of point-to-surface-distance was less for the T1 reconstructions (11.30 mm) than for the Fat reconstructions (16.41 mm).

## 4. Discussion

To the best of our knowledge, there are only two studies [39,40] which analyze the relationship between posture and the muscular system. Pomero et al. [39,40] used a method, which combined stereo-radiographic 3D reconstructions of the spine, forceplate acquisition, physical testing (Cybex II), MRI reconstruction, and muscle modeling. While preliminary results indicate that establishing a mechanical model of the spine is



Table 6

Distance points surfaces in mm. Mean, Std, Min and Max 2xRMS with the average, standard deviation, minimal and maximal of 2xRMS on all reconstructions of one muscle (9 reconstructions/per muscle).

	Projection: distance points surfaces in mm							
	Mean 2xRMS <sup>a</sup>		Std 2xRMS		Min 2xRMS <sup>a</sup>		Max 2xRMS	
	T1	Fat	T1	Fat	T1	Fat	T1	Fat
Mean	2.62	2.95	0.48	0.52	1.96	2.36	3.80	4.30
Std	0.73	0.91	0.35	0.53	0.57	0.63	1.85	2.72
Min	1.27	1.32	0.16	0.09	0.88	1.15	1.83	1.57
Max	4.15	5.80	1.98	2.91	3.24	3.76	11.30	16.41

<sup>a</sup> Significant difference between Fat and T1.

possible, this complex approach does not appear to be clinically applicable, and thus carries limited practical value.

As mechanical models and diagnostic protocols can offer significant contributions to the treatment of degenerative pathologies and deformities, the need exists for further research into the role of soft tissue stabilizers in postural balance.

The reproducibility of the DPSO methods with MR images was previously established by Sudhoff et al. [29] for the muscles involved in knee motion. Reconstructions were based on volume interpolated breath-hold examination (VIBE) images. In order to obtain good visualization of soft tissue, the authors used T1 images. The correlation between the intensity of the signal and the quantification of fat was not reliable due to the lack of homogeneity of the magnetic field.

In order to quantify both muscle geometry and fat infiltration, our study utilized the three point Dixon method. This method permits calculation of water and fat images. The method was originally developed by Dixon [30] using a two point (in phase and out of phase) acquisition. The two point Dixon method is sensitive to inhomogeneity of the magnetic field. To avoid this problem, Glover and Schneider introduced the three point Dixon method [31]. As demonstrated by Bley et al. [41], this method is robust and, as for CT analysis, permits separation of the fat and water volumes (Fat and Water images), thus allowing quantification of fat infiltration.

The time needed to complete the 3D reconstruction of all the studied muscles from the thoraco-lumbar region to the patella (Table 1), with a systematic outline of every slice (reference methods), was between 14 and 15 hours, while the time needed to obtain the reconstruction with Muscl'X software and DPSO methods was about 7 hours. The substantial reduction in time renders the software compatible for clinical research but further research should aim at reducing the time needed for reconstruction.

In this study, the reference object was constructed based on T1 images; this sequence was chosen based on the contrast between fat and muscle and for its wide use in clinical applications. The agreement between the two methods (T1 and Fat images) was variable between muscles and comprised between  $-4.50\%$  and  $+8.00\%$  for the mean AS, the length and the volume. These values of agreement reflect two study volunteers and analysis of 36 muscles. For the same muscle, it appeared that on average the volume and the mean AS obtained with the DPSO method was smaller (T1:  $\sim 1\%$ , Fat:  $\sim 2\%$ ) than the volume obtained with the

reference method. The difference was significantly greater for the DPSO method with Fat image. This systematic bias could be explained by the use of the contrast optimization which can reduce the AS of the non-manually segmented slices. However, this error did not hamper the use of the DPSO method in the 3D geometry assessment of the muscle.

The segmentation of the three point Dixon method had the same quality of segmentation as the segmentation on T1 images. The Fat images had higher fat/muscle contrast than the T1 images; however, visibility of the epimysium and the bone/muscle contrast were better delineated in the T1 images (Fig. 4). This may also explain why the systematic error in the agreement between methods was greater for DPSO with Fat images than for DPSO with T1 images.

An analysis of the accuracy of muscle shape (i.e. comparison of T1/Fat reconstructions with the reference) revealed small point-to-surface-distance errors (mean 2xRMS errors of 2.6 mm and 2.95 mm for T1 and Fat reconstructions, respectively). The references were only constructed on the T1 image, which could explain a greater difference in terms of point to surface on the Fat images. The DPSO method leads to error when a muscle is poorly approximated by an ellipse (e.g. the adductor), or when there are abrupt variations in muscle shape between adjacent slices (e.g. the iliacus). However, errors in terms of shape had little impact on the agreement for other parameters: muscle length, mean AS, and volumes between the two methods. The greatest impact was found on measurement of the maximal AS for which the limits of agreement was  $[-9\%; 11\%]$  for T1 and  $[-11\%; 13.50\%]$  for Fat.

Point-to-surface errors from this study were comparable to previously published values [27–29]. Sudhoff [29] reported 6% error in muscle volume when compared to a reference object (based on 10 volunteers and 2 operators). Using CT images, Jolivet et al. [28] reported errors (two standard deviations around the mean volume) of 5–12% in a reproducibility analysis of the hip muscles of 30 subjects (3 operators). In the current study, which includes only two volunteers, errors were on average, less than 5% for all parameters. Of note, the current protocol used pre-defined slices to identify the muscle insertion.

The variation of reproducibility among the different muscles may be explained by several reasons. First, independent of the image considered, the distinction between muscles was not always straightforward (e.g. gluteus minimus versus gluteus medius or vastus medialis versus vastus intermedius)—a

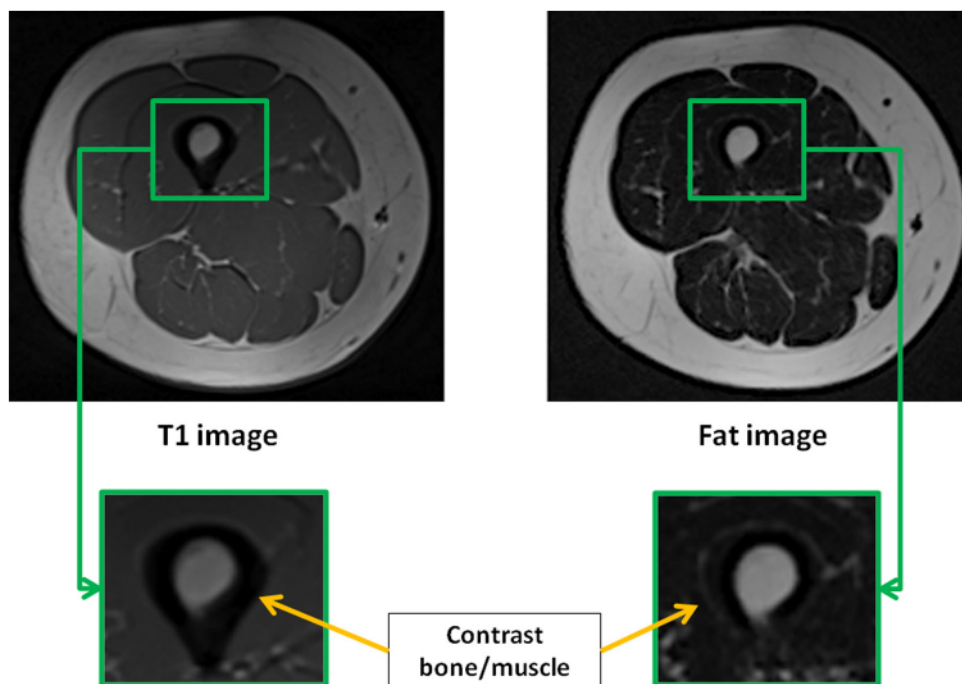


Fig. 4. Contrast between bone/muscle in T1 and Fat images.

difficulty previously reported by other authors [29]. Second, MRI acquisitions were acquired in free breathing, which led to motion artifacts in the abdominal region, that may compromise accuracy of segmentation of the abdominal muscles (e.g. the rectus abdominus and obliques).

In conclusion, this study presents a validation of a protocol to characterize muscle geometry and quantification of the fat infiltration with MRI. The combination of the MuscI'X software with the DPSO method and the three point Dixon method demonstrated good agreement, a reproducibility of less than 5% and led to a substantial gain in reconstruction time. The imaging protocol included in this work is broadly available for clinical scans and has the potential to assess muscular differences between patients and thus the ability to generate patient-specific musculoskeletal models.

## References

- [1] Cho KJ, Suk SI, Park SR, et al. Risk factors of sagittal decompensation after long posterior instrumentation and fusion for degenerative lumbar scoliosis. *Spine (Phila Pa 1976)* 2010;35:1595–601.
- [2] Glassman SD, Berven S, Bridwell K, et al. Correlation of radiographic parameters and clinical symptoms in adult scoliosis. *Spine (Phila Pa 1976)* 2005;30:682–8.
- [3] Glassman SD, Bridwell K, Dimar JR, et al. The impact of positive sagittal balance in adult spinal deformity. *Spine (Phila Pa 1976)* 2005;30:2024–9.
- [4] Lafage V, Schwab F, Patel A, et al. Pelvic tilt and truncal inclination: two key radiographic parameters in the setting of adults with spinal deformity. *Spine (Phila Pa 1976)* 2009;34:E599–606.
- [5] Gille O, Jolivet E, Dousset V, et al. Erector spinae muscle changes on magnetic resonance imaging following lumbar surgery through a posterior approach. *Spine (Phila Pa 1976)* 2007;32:1236–41.
- [6] Kawaguchi Y, Matsui H, Tsuji H. Back muscle injury after posterior lumbar spine surgery. Part 2: histologic and histochemical analyses in humans. *Spine* 1994;19:2598–602.
- [7] Kawaguchi Y, Matsui H, Tsuji H. Back muscle injury after posterior lumbar spine surgery. A histologic and enzymatic analysis. *Spine* 1996;21:941–4.
- [8] Taylor H, McGregor AH, Medhi-Zadeh S, et al. The impact of self-retaining retractors on the paraspinal muscles during posterior spinal surgery. *Spine* 2002;27:2758–62.
- [9] Weber BR, Grob D, Dvorak J, et al. Posterior surgical approach to the lumbar spine and its effect on the multifidus muscle. *Spine* 1997;22:1765–72.
- [10] Flicker PL, Fleckenstein JL, Ferry K, et al. Lumbar muscle usage in chronic low back pain. Magnetic resonance image evaluation. *Spine* 1993;18:582–6.
- [11] Gejo R, Matsui H, Kawaguchi Y, et al. Serial changes in trunk muscle performance after posterior lumbar surgery. *Spine* 1999;24:1023–8.
- [12] Kim DY, Lee SH, Chung SK, et al. Comparison of multifidus muscle atrophy and trunk extension muscle strength: percutaneous versus open pedicle screw fixation. *Spine* 2005;30:123–9.
- [13] Humphrey AR, Nargol AV, Jones AP, et al. The value of electromyography of the lumbar paraspinal muscles in discriminating between chronic-low-back-pain sufferers and normal subjects. *Eur Spine J* 2005;14:175–84.
- [14] Mooney V, Gulick J, Perlman M, et al. Relationships between myoelectric activity, strength, and MRI of lumbar extensor muscles in back pain patients and normal subjects. *J Spinal Disord* 1997;10:348–56.
- [15] Barker KL, Shamley DR, Jackson D. Changes in the cross-sectional area of multifidus and psoas in patients with unilateral back pain: the relationship to pain and disability. *Spine* 2004;29:E515–9.
- [16] Dangaria TR, Naesh O. Changes in cross-sectional area of psoas major muscle in unilateral sciatica caused by disc herniation. *Spine* 1998;23:928–31.
- [17] Danneels LA, Vanderstraeten GG, Cambier DC, et al. CT imaging of trunk muscles in chronic low back pain patients and healthy control subjects. *Eur Spine J* 2000;9:266–72.
- [18] Hultman G, Nordin M, Saraste H, et al. Body composition, endurance, strength, cross-sectional area, and density of MM erector spinae in men with and without low back pain. *J Spinal Disord* 1993;6:114–23.
- [19] Keller A, Gunderson R, Reikeras O, et al. Reliability of computed tomography measurements of paraspinal muscle cross-sectional area and density in patients with chronic low back pain. *Spine* 2003;28:1455–60.
- [20] Mayer TG, Vanharanta H, Gatchel RJ, et al. Comparison of CT scan muscle measurements and isokinetic trunk strength in postoperative patients. *Spine* 1989;14:33–6.

- [21] Parkkola R, Rytokoski U, Kormano M. Magnetic resonance imaging of the discs and trunk muscles in patients with chronic low back pain and healthy control subjects. *Spine* 1993;18:830–6.
- [22] Savage RA, Millerchip R, Whitehouse GH, et al. Lumbar muscularity and its relationship with age, occupation and low back pain. *Eur J Appl Physiol Occup Physiol* 1991;63:265–8.
- [23] Storheim K, Holm I, Gunderson R, et al. The effect of comprehensive group training on cross-sectional area, density, and strength of paraspinal muscles in patients sick-listed for subacute low back pain. *J Spinal Disord Tech* 2003;16:271–9.
- [24] Airaksinen O, Herno A, Kaukanen E, et al. Density of lumbar muscles 4 years after decompressive spinal surgery. *Eur Spine J* 1996;5:193–7.
- [25] Salminen JJ, Erkintalo-Tertti MO, Paajanen HE. Magnetic resonance imaging findings of lumbar spine in the young: correlation with leisure time physical activity, spinal mobility, and trunk muscle strength in 15-year-old pupils with or without low-back pain. *J Spinal Disord* 1993;6:386–91.
- [26] Tracy BL, Ivey FM, Jeffrey Metter E, et al. A more efficient magnetic resonance imaging-based strategy for measuring quadriceps muscle volume. *Med Sci Sports Exerc* 2003;35:425–33.
- [27] Jolivet E, Daguet E, Pomero V, et al. Volumic patient-specific reconstruction of muscular system based on a reduced dataset of medical images. *Comput Methods Biomech Biomed Engin* 2008;11:281–90.
- [28] Jolivet E [PhD report] Biomechanical modelisation of hip and its soft tissue for hip fracture risk. Paris: Laboratory of Biomechanics. ENSAM; 2007. p. 177.
- [29] Sudhoff I, de Guise JA, Nordez A, et al. 3D-patient-specific geometry of the muscles involved in knee motion from selected MRI images. *Med Biol Eng Comput* 2009;47:579–87.
- [30] Dixon WT. Simple proton spectroscopic imaging. *Radiology* 1984;153:189–94.
- [31] Glover GH, Schneider E. Three-point Dixon technique for true water/fat decomposition with B0 inhomogeneity correction. *Magn Reson Med* 1991;18:371–83.
- [32] Glover GH. Multipoint Dixon technique for water and fat proton and susceptibility imaging. *J Magn Reson Imaging* 1991;1:521–30.
- [33] Yeung HN, Kormos DW. Separation of true fat and water images by correcting magnetic field inhomogeneity in situ. *Radiology* 1986;159:783–6.
- [34] Angotti A. Contribution à l'analyse musculaire de patients présentant des pathologies rachidiennes. Paris: Laboratoire de Biomecanique. ENSAM; 2008. p. 55.
- [35] Tochu F. A contouring program based on dual kriging interpolation. *Eng Comput* 1993;0:160–77.
- [36] Martin Bland J. Statistical methods for assessing agreement between two methods of clinical measurement. *Lancet* 1986;307–10.
- [37] Standardization II Of. Accuracy (trueness and precision) of measurement methods and results – Part 2: basic method for the determination of repeatability and reproducibility of a standard measurement method: 1994, 1994.
- [38] Mitton D, Landry C, Veron S, et al. 3D reconstruction method from biplanar radiography using non-stereo corresponding points and elastic deformable meshes. *Med Biol Eng Comput* 2000;38:133–9.
- [39] Pomero V, Lavaste F, Imbert G, et al. A proprioception based regulation model to estimate the trunk muscle forces. *Comput Methods Biomech Biomed Engin* 2004;7:331–8.
- [40] Pomero V, Vital JM, Lavaste F, et al. Muscular modelling: relationship between postural default and spine overloading. *Stud Health Technol Inform* 2002;88:321–5.
- [41] Bley TA, Wieben O, Francois CJ, et al. Fat and water magnetic resonance imaging. *J Magn Reson Imaging* 2010;31:4–18.